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Introduction

Valence tautomeric (VT) metal complexes are well known systems showing molecular bistability.^{1,2} The close energies of metal- and ligand-based redox-active orbitals allow a reversible intramolecular electron transfer to occur, which gives rise to distinctly different electronic states (redox isomers). Particularly interesting are VT cobalt dioxolenes because intramolecular electron transfer is accompanied by a spin state change at the cobalt center resulting in reversible switching between diamagnetic low-spin (ls) cobalt(iii) $(S = 0)$ and paramagnetic high-spin (hs) cobalt(II) $(S = 3/2)$ ions. Greatly differing magnetic properties of the two states and the opportunity to switch reversibly between them by external stimuli render VT cobalt complexes highly attractive for applications as molecular switches, sensors and molecule-based memory units.³⁻⁶

The switching in VT metal complexes can be triggered by temperature, pressure, magnetic field, soft X-rays or by light.⁷

Most of the research in the field is dedicated to the development of thermally switchable VT species. However, switching with light is much more attractive for prospective applications due to high speed of addressing, superior resolution, and high selectivity. It is known that photoswitching in VT metal complexes can be achieved by irradiation into appropriate charge transfer (CT) absorption bands.⁸ However, the lifetime of such photoinduced metastable states is in the order of nanoseconds at room temperature (RT). The metastable states can be stabilized by cooling which allows the photoswitching to be performed only at very low temperatures, usually below 20 K.⁹⁻¹² Shultz et al. were able to stabilize photoinduced states at higher temperatures up to 90 K using crystal lattice effects.13,14 However, the photoswitching becomes impossible at molecular level in such a case. Thus, all previously reported VT photoswitches operate only at very low temperatures, which results in serious limitations for their use in molecular devices.

below 20 K) because photoinduced states are highly unstable at room temperature. The thermal instability hinders any practical application of these complexes in genuine devices. In this report, for the first time we demonstrate photoswitching of VT species and associated magnetic properties at room temperature. The bidirectional photoswitching in solution is due to cis-trans photoisomerizable 4-styrylpyridine ligands deliberately integrated into cobalt dioxolene molecular complexes. The novel type of photoswitching has been coined Ligand-Driven Light-Induced Valence Tautomerism (LD-LIVT). The photoconversion of VT states of 28% has been achieved in solution at room temperature. The photoinduced states show extraordinary thermal stability for hours at room temperature, as compared to common nanoseconds reported previously. The switching proceeds at molecular level with the effective photoswitching rate of 3×10^{13} molecules per s under our conditions. Consequently, this work may open new horizons in applications of molecular switches based on VT metal complexes in molecular EDGE ARTICLE

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Valence tautomeric (VT) metal complexes are highly promising bistable molecular compounds for applications as molecular switches in molecular electronics and spintronics. Although VT species can be switched with light, the photoswitching in all reported systems requires very low temperatures (usually

Bidirectional photoswitching of magnetic

light-induced valence tautomerism†

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properties at room temperature: ligand-driven

devices functioning at room temperature.

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[†] Electronic supplementary information (ESI) available: Derivation of the non-linear regression fitting functions, crystallographic details, magnetization and electrochemical data, EPR and electronic absorption spectra. CCDC 1015850–1015852 and 1058866. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c5sc00130g

Nonetheless, a photoinduced state might be stabilized by introducing photoisomerizable ligands into VT systems¹⁵ in such a way that a ligand-based photoreaction affects the energy of redox-active orbitals and thus triggers intramolecular electron transfer. The stability of such photoinduced state is dictated by the stability of the photoisomer of the ligand which can exceed years at RT.¹⁶ The very first step in this direction has been attempted recently by Frank et al .¹⁷ Although their spirooxazine-derived ligand maintained photoactivity within a VT cobalt complex, no solid evidence for the photoswitching of VT states at RT has been provided. Thus, VT metal complexes that can be switched with light at RT remained unknown to date.

Our group works on stabilization of photoinduced states in VT and spin-crossover metal complexes by integration of photoisomerizable ligands into bistable species.¹⁸–²⁰ Very recently, we have developed a VT cobalt dioxolene system featuring two photoactive trans-4-phenylazopyridine ligands and introduced a new concept for switching VT states called "Coordination-Induced Valence Tautomerism" (CIVT).²⁰ Unfortunately, low thermal stability of photogenerated cis-4-phenylazopyridine precluded a detailed examination of photoswitching in this system. To improve photophysical properties of the system we decided to substitute 4-phenylazopyridine with 4-styrylpyridine (4-stypy) ligands that show high thermal stability of both cisand trans-isomers.²¹ Chemical Science

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Here, we report two VT cobalt complexes *trans*-6 and *cis*-6 containing photoactive trans-4-stypy and cis-4-stypy ligands, respectively (Scheme 1). Their molecular and electronic structures are thoroughly investigated by variable-temperature X-ray crystallography, magnetic susceptibility measurements, NMR, EPR and electronic absorption spectroscopy, electrochemistry and titration experiments. For the first time, the electronic states of VT metal complexes are switched with light at room temperature. Consequently, the switching of VT states allows to

Scheme 1 Cobalt dioxolene complexes trans-5 and trans-6 with their respective electronic structures: $[Is-Co^{III}(Cat)(SO)(trans-4-stypy)₂]$ (trans- 6^{LS}), [hs-Co^{II}(SQ)₂(trans-4-stypy)₂] (trans- 6^{HS}), and [hs-Co^{II}(SQ)₂(trans-4-stypy)] (trans-5). Molecular and electronic structures of cis-5 and cis-6 are similar except of trans-4-stypy substituted by cis-4-stypy.

control magnetic properties with light at ambient temperatures. This opens new horizons for application of molecular switches based on VT metal complexes in functioning molecular devices.

Experimental section

Materials

All starting materials and solvents were used as received without further purification unless otherwise noted. Pure anhydrous solvents, required for work under inert atmosphere, were collected from a solid state solvent purification system (Glass Contour System, Irvine, CA) and stored over activated molecular sieves.

Syntheses

Precursor cobalt tetramer was obtained as a benzene solvate $[Co(tbdiox)₂]$ ₄ (C_6H_6) _{2.75} by a synthetic procedure described by Pierpont et al.²² trans-4-Styrylpyridine (trans-4-stypy) was prepared according to a method described by Chiang and Hartung.²³ cis-4-Styrylpyridine (cis-4-stypy) was obtained in a Wittig reaction as reported by Williams et $al.^{21}$.

 $[Co(tbdiox)₂(trans-4-stypy)₂]$ (trans-6) was prepared according to a common procedure for bis(o-dioxolene) cobalt complexes:¹ under inert atmosphere, trans-4-styrylpyridine (200 mg, 1.10 mmol) dissolved in toluene (10 mL) was added dropwise to $[Co(tbdiox)₂]$ ₄ (C₆H₆)_{2.75} (300 mg, 0.14 mmol) dissolved in hot toluene (50 mL, 80 $^{\circ}$ C). The reaction mixture was stirred overnight at 80 °C and then stored at -35 °C for 3 days before a greygreen precipitate was collected and dried in vacuo (274 mg, yield: 58%). Elemental analysis calcd (%) for $C_{54}H_{62}CoN_2O_4$: C 75.24, H 7.25, N 3.25; found: C 74.99, H 7.24, N 3.32.

 $[Co(tbdiox)₂(cis-4-stypy)₂]$ (cis-6) was obtained following a similar synthetic protocol: under inert atmosphere, cis-4-styrylpyridine (295 mg, 1.63 mmol) was added to a hot toluene solution (70 mL, 80 °C) of $[Co(tbdiox)_2]_4 \cdot (C_6H_6)_{2.75}$ (448 mg, 0.20 mmol). The reaction mixture was stirred for 2 days at 80 $^{\circ}$ C and then stored at -35 °C for several days before a black crystalline precipitate was collected and dried in vacuo (503 mg, yield: 73%). Elemental analysis calcd (%) for $C_{54}H_{62}CoN_2O_4$: C 75.24, H 7.25, N 3.25; found: C 75.83, H 7.13, N 3.36.

Instrumentation and physical measurements

Elemental analyses were carried out with a EURO EA analyser from Euro Vector. Magnetic susceptibility data on solid samples were collected using a Quantum Design MPMS-XL SQUID magnetometer. The data were obtained for microcrystalline samples restrained within a polycarbonate gel capsule. DC susceptibility data were collected in the temperature range 2–400 K at applied magnetic field of 1 T. The program Ju X was used for the simulation and analysis of magnetic data.²⁴ Electrochemical measurements were performed under nitrogen atmosphere at RT using a standard three-electrode setup with glassy carbon working electrode and platinum rods as counter and reference electrodes. The potentiostat was a μ Autolab Type-III. Analyte solutions were prepared in CH_2Cl_2 containing 0.1 M $n_{\text{Bu}_4\text{NPF}_6}$ as supporting electrolyte. All potentials are referenced

to the $\mathrm{Fc}^{+/0}$ redox couple measured after adding ferrocene to the analyte solution. EPR spectra were recorded on a Jeol CW spectrometer JES-FA200 equipped with an X-band Gunn diode oscillator bridge and a cylindrical mode cavity. Simulations were performed using the program W95EPR written by F. Neese.²⁵ Irradiation experiments were conducted in situ through a quartz window in the cavity. NMR spectra of trans-6 in solution were recorded in rotating 5 mm o.d. tubes with Jeol JNM-LA 400 FT NMR spectrometer and processed with Delta V4.0 software provided by Jeol Ltd. NMR spectra of cis-6 in solution were obtained without rotating, in 5 mm o.d. tubes on a Bruker Avance DRX 400 WB spectrometer and processed with TopSpin 1.3 software. Magnetic susceptibility in solutions were determined by the Evans NMR method.²⁶ During the variable temperature and titration experiments the outer tube (standard 5 mm o.d. NMR tube with a PTFE spindle valve) contained a reference solvent mixture $C_7H_8/C_7D_8/TMS$ (10/2/1), while the inner tube (capillary tube with 1.5 mm diameter sealed with inert wax) contained the paramagnetic complex in the same solvent mixture. The irradiation experiments were performed using a rotating 5 mm o.d. quartz NMR tube with a PTFE spindle valve with the complex solution in the outer tube and the reference solvent mixture in the inner capillary tube. Electronic absorption spectra were recorded with a Shimadzu UV 3600 spectrophotometer. The samples were prepared under anaerobic conditions and sealed in QS Quartz Suprasil cells (10 mm light path) with PTFE spindle valves. Variable-temperature spectra were recorded with an Analytik Jena SPECORD S600 spectrophotometer. The samples were prepared under anaerobic conditions and measurements were conducted inside a glove box using QS Quartz Suprasil cells (10 mm light path). The solutions were continuously stirred with a magnet bar and Edge Article

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the temperature inside the cells was monitored. An LOT-Oriel $Xe(OF)$ arc lamp (1 kW) equipped with an Omni- λ 300 monochromator was used as a wavelength-variable light source in all irradiation experiments except the in situ irradiation followed by EPR spectroscopy. For the latter, an LOT-Oriel Xe(OF) arc lamp (150 W) equipped with an Andover bandpass filter (CWL: 322.9 nm, Tmax: 28.0%, FWHM: 10.6 nm) were employed.

X-ray crystallographic data collection and structure refinement

Suitable crystals were embedded in protective per fluoropolyalkyl ether oil and transferred to the cold nitrogen gas stream of the diffractometer. Intensity data for *trans-*6 were collected at 120, 295 and 305 K, and for cis-6 at 100 K on a Bruker Kappa Smart APEX2 (Mo K_{α} radiation, $\lambda = 0.71073$ Å, graphite monochromator). Data were corrected for Lorentz and polarization effects; semi empirical absorption corrections were applied on the basis of multiple scans using SADABS.²⁷ The structures were solved by direct methods and refined by fullmatrix least squares procedures on F^2 using *SHELXTL NT* 6.12.²⁸ All non-hydrogen atoms were refined with anisotropic displacement parameters. Hydrogen atoms were placed in position of optimized geometry and their isotropic displacement parameters were tied to those of their corresponding carrier atoms by a factor of 1.2 or 1.5. SIMU and ISOR restraints were applied in the refinement of the disorder. The crystallographic data, data collection and structure refinement details are summarized in Table 1 (ESI†).

At temperatures of 120 K and 295 K trans-6 was situated on a general position of space group $\overline{P1}$. While at 120 K the molecule was well ordered, disorder was observed at higher temperatures. At 295 K the two pyridyl ligands were subjected to orientational

trans-6 trans-6 trans-6 trans-6 cis-6 \cdot 4(C₇H₈) Temperature, K 120 100 295 305 100 Chemical formula $C_{54}H_{62}CoN_2O_4$ $C_{54}H_{62}CoN_2O_4$ $C_{54}H_{62}CoN_2O_4$ $C_{54}H_{62}CoN_2O_4$ $C_{54}H_{62}CoN_2O_4$ $C_{54}H_{62}CoN_2O_4$ $C_{54}H_{62}CoN_2O_4$ $C_{54}H_{62}CoN_2O_4$ $F_{\rm w}$ 861.99 861.99 861.99 861.99 861.99 861.99 1230.52 Cryst size, mm $0.30 \times 0.26 \times 0.16$ $0.30 \times 0.26 \times 0.16$ $0.30 \times 0.26 \times 0.16$ $0.50 \times 0.34 \times 0.24$ Cryst sys Triclinic Triclinic Triclinic Monoclinic Space group $P\bar{1}$ $P\bar{1}$ $P\bar{1}$ $P\bar{1}$ $P\bar{1}$ $P\bar{2}1/n$ a, \AA 10.6748(4) 10.7956(11) 9.8997(9) 16.5308(6) b, \AA 11.9906(5) 12.0991(13) 10.8120(10) 11.7257(4) c, \AA 19.6875(8) 19.834(2) 12.1278(11) 18.9821(7) α , deg 101.316(2) 102.162(6) 104.002(5) 90
 β deg 101.002(5) 101.0522(17) 101.146(5) 102.270(5) 102.270(5) β , deg 101.9532(17) 101.446(5) 102.270(5) 107.445(2) γ , deg 103.7626(17) 104.036(5) 101.408(5) 90
 $V \stackrel{\delta}{\lambda}$ 1187.40(10) 2313.24(16) 2370.3(4) 1187.40(10) 25 V, \hat{A}^3 2313.24(16) 2370.3(4) 1187.49(19) 3510.2(2) Z 2212 r_{calcd} g cm⁻³ 1.238 1.208 1.205 1.205 1.205 1.164
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Table 1 Crystallographic data, data collection and structure refinement details for trans-6 and cis-6

disorder affecting the whole ligand moieties. Two alternative orientations of the pyridyl ligand featuring inverted orientations of the central $C=C$ double bond were refined resulting in site occupancies of 85.4(3) and 14.6(3)% of the 50 involved atoms (for a graphical representation of the disorder see ESI†). Furthermore, one of the four ^tBu groups was disordered. Here, also two different orientations were refined giving site occupancies of $52(1)$ and $48(1)\%$ for the atoms C22–C24 and C22A–C24A, respectively.

The situation changed at temperatures above 295 K. At 305 K the complex molecule was now located on a crystallographic inversion center and exhibited crystallographically imposed C_i symmetry. As in the 295 K structure the pyridyl ligands were subjected to disorder with different orientations of the central $C=C$ double bond and were refined resulting in site occupancies of 83.6(6) and 16.4(6)% of the involved atoms. Furthermore, one of the crystallographically independent 'Bu groups was disordered. Here, again two different orientations were refined giving site occupancies of $69(2)$ and $31(2)\%$ for the atoms C8–C10 and C8A–C10A, respectively. All three crystal structures of trans-6 point to a ls-Co^{III}(Cat)(SQ) electronic configuration (vide infra). However, owing to C_i symmetry at 305 K one cannot distinguish between a mixed-valent delocalized ls- $Co^{III}(Cat)(SQ)$ \leftrightarrow ls-Co^{III}(SQ)(Cat) electronic structure and a positional disorder in the localized ls-Co^{III}(Cat)(SQ) form.^{14,20} The observation of a crystallographic inversion center at higher temperatures may be due to increasing thermal vibration of atoms accompanied by the incipient valence tautomeric transition as confirmed by magnetic measurements. Chemical Science

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Molecules of cis-6 at 100 K are located on a crystallographic inversion center and exhibit C_i symmetry. The compound crystallized with four molecules of toluene per formula unit. One of the two independent toluene molecules was disordered. Two alternative orientations were refined resulting in site occupancies of 61(1) and 39(1)% for the atoms C201–C207 and C211–C217. SAME and SIMU restraints were applied in the refinement of the disordered toluene.

Results and discussion

Syntheses and crystal structures

The neutral complexes trans-6 and cis-6 were synthesized according to a common procedure for VT cobalt dioxolenes.¹ Slow precipitation from toluene solutions afforded crystals suitable for X-ray structure determination.²⁹ At 120 K trans-6 reveals a distorted octahedral geometry featuring two equatorial bidentate *o*-dioxolene and two axial monodentate *trans*-4-stypy ligands (Fig. 1). Short Co–O $(1.854(1)\cdots1.916(1)$ Å) and Co–N $(1.938(1)$ and $1.945(1)$ Å) distances are characteristic for a lowspin (ls) cobalt (m) ion. Intraligand C–O and (O) C–C (O) bond lengths, generally diagnostic of the oxidation level of o-dioxolenes,³⁰ indicate the presence of a cate cholate dianion (Cat²⁻) and a semi-quinone monoanion π -radical (SQ⁻). The SQ state for one of the ligands is further confirmed by quinoid-type distortion with alternating long-short bonds. Thus, the ligand mixed valency is localized in the solid state at low temperatures.

Fig. 1 Molecular structure of trans-6 determined at 120 K. Thermal ellipsoids are drawn at 50% probability. Hydrogen atoms are omitted for clarity

Fig. 2 Molecular structure of cis-6 determined at 100 K. Thermal ellipsoids are drawn at 50% probability. Hydrogen atoms are omitted for clarity

At $T = 305$ K all cobalt-ligand bonds remain short, but the quinoid-type distortion is observed in both dioxolenes.

The structurally related cis-6 features two axial cis-4-stypy ligands (Fig. 2). Due to steric factors the pyridyl and phenyl rings of cis-4-stypy cannot be coplanar. Thus, the two rings form a dihedral angle of 50.1° confining a twisted ethylene bridge. Similar twisted geometry has been observed previously.^{31,32} Short Co–N $(1.940(1))$ and Co–O $(1.885(1)$ and $1.878(1)$ Å) distances point to a ls-Co^{III} center at 100 K. Similar to trans-6 at 305 K, intraligand bonds of both o -dioxolenes in *cis*-6 are equally subjected to quinoid type distortion corroborating Cat^{2-}/SQ^- ligand mixed valency.

Magnetic properties in solid state

The ls-Co^{III}(Cat)(SQ) ground state for *trans*-6 is corroborated by magnetic susceptibility measurements on a microcrystalline sample revealing an effective magnetic moment $\mu_{\text{eff}} = 1.79 \mu_{\text{B}}$ invariant in the temperature range 8–250 K (see ESI†). The data could be fitted for a spin doublet $S = 1/2$ system to afford $g = 2.064$ and a Weiss constant $\theta = -0.15$ K. Hence, based on both, bond length analysis and magnetic measurements, the electronic structure of solid trans-6 below 250 K is unambiguously assigned as $[Is\text{-}Co^{III}(Cat)(SQ)(trans-4-stypy)_2]$ (trans-6^{LS}). At temperatures above 250 K magnetic moment gradually increased to $4.00\mu_B$ at 400 K revealing a thermally induced trans- 6^{LS} \rightarrow [hs-Co^{II}(SQ)₂(trans-4-stypy)₂] (trans-6^{HS}; hs = high-spin) transition. The data were fitted with the van't Hoff equation (see ESI†): with the high-temperature limit of magnetic moment fixed at $5.0\mu_B$ as a common value for hs-Co^{II}(SQ)₂ isomers,¹³ our best fit provides an enthalpy change $\Delta H = 41(1)$ kJ mol⁻¹ and an entropy change $\Delta S = 104(3)$ J mol⁻¹ K⁻¹.³³

Similar to trans-6, magnetic data for solid cis-6 below 250 K were fitted for an $S = 1/2$ spin system with $g = 2.073$ and θ = -0.62 K (see ESI†); and the electronic structure was assigned as [ls-Co^{III}(Cat)(SQ)(cis-4-stypy)₂] (cis- 6^{LS}). The thermal transition $cis\text{-}6^{\text{LS}}$ \rightarrow [hs-Co^{II}(SQ)₂(cis-4-stypy)₂] (cis-6^{HS}), indicated by the increase in μ_{eff} at higher temperatures, was fitted to yield $\Delta H = 52(2)$ kJ mol⁻¹ and $\Delta S = 131(3)$ J mol⁻¹ K⁻¹ (see ESI†). The thermodynamic parameters obtained for both trans-6 and cis-6 in solid state are in agreement with typical thermodynamic parameters for thermally induced VT transitions.³⁴ The evolution of μ_{eff} with temperature was fully reversible without detectable hysteresis in both cases. Transition temperatures $T_{1/2}$ in solid state were estimated at 394 and 397 K for trans-6 and cis-6, respectively. Edge Article

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Electrochemistry

Cyclic voltammograms obtained for a $CH₂Cl₂$ solution of trans-6 at RT display one oxidation wave with a half-wave potential $E_{1/2}$ = -0.33 V and two reduction events at $E_{1/2} = -0.67$ and -1.12 V versus $Fc^{+/0}$ (see ESI†). The ratio between normalized oxidative and reductive peak currents for all three waves are near unity pointing to reversible processes. However, the peak separations ΔE for reduction events significantly exceed the ideal value of 58 mV expected for a diffusion-controlled reversible one-electron transfer. Very similar cyclic voltammograms measured for cis-6 show one oxidation at $E_{1/2} = -0.29$ V and two reductions at $E_{1/2} = -0.69$ and -1.11 V (see ESI†). All observed redox events were assigned to the redox-active bis(dioxolene)cobalt core. Yet, a detailed assignment is difficult due to partial dissociation of the complexes in solution (vide infra).

Magnetic properties in solution

The effective magnetic moment of *trans*-6 in solution determined by the Evans NMR method²⁶ remained nearly constant at $1.7\mu_B$ below 215 K, which is in agreement with a pure $\text{ls-Co}^{\text{III}}(\text{Cat})(\text{SQ})$ state. With rising temperature, μ_{eff} gradually increased reaching 4.91(5) μ _B at 353 K (Fig. 3). These data were fitted using the van't Hoff equation to yield $\Delta H = 43(5)$ kJ mol⁻¹, $\Delta S = 144(16)$ J mol^{-1} K⁻¹ and the high-temperature limit of magnetic moment of 5.1(1) μ_B that is characteristic for a pure hs-Co^{II}(SQ)₂ state. The transition temperature $T_{1/2}$ was estimated at 299 K. At RT this solution contains 44% of hs-Co^{II}(SQ)₂ and 56% of $\text{ls-Co}^{\text{III}}(\text{Cat})(\text{SQ})$ species as estimated from RT magnetic moment of 3.59(6) $\mu_{\rm B}$.

Fig. 3 Temperature dependent effective magnetic moment of trans-6 dissolved in toluene determined by the Evans method (toluene/ $[D_8]$ toluene/TMS = $10 : 2 : 1$). Van't Hoff fit parameters: $\Delta H = 43(5)$ kJ mol⁻¹, $\Delta S = 144(16)$ J mol⁻¹ K⁻¹; low- and high-temperature magnetic moments: $\mu_{\text{eff}}(\text{LT}) = 1.69(3)\mu_{\text{B}}$ and $\mu_{\text{eff}}(\text{HT}) = 5.1(1)\mu_{\text{B}}$. Estimated transition temperature: $T_{1/2} = 299$ K.

Fig. 4 Temperature dependent effective magnetic moment of cis-6 dissolved in toluene determined by the Evans method (toluene/ $[D_8]$ toluene/TMS = $10 : 2 : 1$). Van't Hoff fit parameters: $\Delta H = 67(9)$ kJ mol⁻¹, $\Delta S = 232(33)$ J mol⁻¹ K⁻¹; low- and high-temperature magnetic moments: $\mu_{eff}(LT) = 2.45(3)\mu_B$ and $\mu_{eff}(HT) = 4.44(3)\mu_B$. Estimated transition temperature: $T_{1/2} = 287$ K.

Magnetic moment of cis-6 in solution changes gradually from 2.42(6) μ_B at 218 K to 4.52(6) μ_B at 348 K (Fig. 4). The data was fitted to give $\Delta H = 67(9)$ kJ mol⁻¹ and $\Delta S = 232(33)$ J mol⁻¹ K^{-1} . The high temperature limit for magnetic moment is 4.44(3) μ_B that is within the range of typical values for a pure hs- $Co^H(SQ)₂$ state.³⁵ However, the low temperature limit of 2.45(3) μ_B is higher than expected for a pure ls-Co^{III}(Cat)(SQ) state. This might be due to the presence of some paramagnetic impurities in solution of *cis*-6.²⁰ The transition temperature $T_{1/2}$ was estimated at 287 K. Note, that the $T_{1/2}$ value for cis-6 is slightly lower than that for trans-6 in solution (Table 2).

Electronic absorption spectra in solution

The RT electronic absorption spectrum of trans-6 dissolved in toluene is dominated by a strong band at 307 nm ($\varepsilon = 6.1 \times 10^4$) M^{-1} cm⁻¹) arising from $\pi \to \pi^*$ transitions of *trans*-4-stypy,

which overlap with intraligand dioxolene bands (see ESI \dagger).^{36,37} A relatively weak broad band at 750 nm $(\epsilon = 1.8 \times 10^3 \,\mathrm{M}^{-1} \,\mathrm{cm}^{-1})$
with a shoulder at ϵ 600 nm are shared existing CT transitions in with a shoulder at \sim 600 nm are characteristic CT transitions in VT cobalt dioxolenes.² The broad band at \sim 2650 nm, blighted by solvent overtones in the short-wavelength infrared (SWIR) region, was assigned as an intervalence ligand-to-ligand charge transfer (IVLLCT) transition, which is a spectral fingerprint of mixed-valent ls-Co^{III}(Cat)(SQ) species (Fig. 7).³⁵ The half width $\Delta \nu_{\text{exp}}$ of this band was estimated at 860 cm⁻¹. This value is
emailer than the helf width coloulated using the Hugh cquation smaller than the half width calculated using the Hush equation $\Delta v_{\text{calcd}} = (2310 \times v_{\text{max}})^{1/2} = 2981 \text{ cm}^{-1}$ ³⁸ Consequently, *trans-*
 ϵ^{15} behaves as a class III fully delegating lixed mixed valent 6LS behaves as a class-III fully delocalized ligand mixed-valent system in solution.

At 320 K the solution of trans-6 appeared green colored and the 600 nm absorption was nearly buried beneath the broad 750 nm band. Upon cooling to 259 K, the solution turned bluishgreen, the 750 nm band decreased in intensity, and the absorption at 600 nm became more prominent (Fig. 5), which indicates a hs-Co^{II}(SQ)₂ \rightarrow ls-Co^{III}(Cat)(SQ) transition.²⁰ Since the available region of our spectrophotometer for variabletemperature measurements is limited to 1020 nm, the spectral evolution in the SWIR region could not be recorded. The temperature dependence of the 750 nm band was fitted according to the van't Hoff equation to yield $\Delta H = 56(1)$ kJ mol⁻¹ and $\Delta S = 190(5)$ J mol⁻¹ K⁻¹ (Fig. 6). Consequently, the estimated $T_{1/2}$ value for *trans*-6 is 295 K that is in very good agreement with the value determined by NMR spectroscopy (299 K).

Fig. 5 Temperature-dependent electronic absorption spectra of trans-6 dissolved in toluene.

Fig. 6 Temperature dependence of the 750 nm absorption band of trans-6 and cis-6 dissolved in toluene. van't Hoff fit parameters for trans-6: $\Delta H = 56(1)$ kJ mol⁻¹, $\Delta S = 190(5)$ J mol⁻¹ K⁻¹; low- and hightemperature molar extinction coefficients: ε (LT) = 0.908(3) \times 10³ M⁻¹ cm⁻¹ and ε (HT) = 2.490(4) \times 10³ M⁻¹ cm⁻¹, estimated transition
temperature T₁₁ - 295 K. Eit parameters for cis-6; AH - 52(3) k1 mol⁻¹ temperature $T_{1/2} = 295$ K. Fit parameters for cis-6: $\Delta H = 52(3)$ kJ mol⁻¹, $\Delta S = 184(11)$ J mol⁻¹ K⁻¹; low- and high-temperature molar extinction coefficients: ε (LT) = 1.041(9) \times 10³ M⁻¹ cm⁻¹ and ε (HT) = 2.337(5) \times 10³ M^{-1} cm⁻¹, estimated transition temperature $T_{1/2} = 283$ K.

Similar to *trans*-6, a strong band at 306 nm ($\epsilon = 4.2 \times 10^4$) M^{-1} cm⁻¹) caused by *cis*-4-stypy and dioxolene intraligand transitions was observed for cis-6 solution at RT (see ESI†). As compared to *trans*-6, the shoulder at \sim 600 nm is less developed and the IVLLCT transition at \sim 2650 nm is weaker for *cis*-6 (see ESI[†]). These two features indicate that at RT the ls-Co^{III}(Cat)(SQ) fraction in *cis-*6 solution is lower than in *trans-*6 solution. Thus, the VT equilibrium is more shifted toward the hs-Co^{II}(SO)₂ species in case of *cis-*6, which is very likely due to weaker coordination of cis-4-stypy (vide infra).

Similar to trans-complex, a color change was observed upon cooling a *cis-*6 solution. The temperature dependence of absorption at 750 nm was fitted to give $\Delta H = 52(3)$ kJ mol⁻¹ and $\Delta S = 184(11)$ J mol⁻¹ K⁻¹. The estimated $T_{1/2}$ value of 283 K is in very good agreement with the value obtained from the Evans method (287 K) (Fig. 6). Again note, that $T_{1/2}$ values for cis-6 are slightly lower than those of *trans*-6 in solution (Table 2).

Titration with 4-stypy ligands

The thermodynamic parameters obtained for both trans-6 and cis-6 in solution are larger than those in the solid state and

Fig. 7 Changes in absorption spectrum of trans-6 dissolved in toluene upon titration with trans-4-stypy at RT. Signals marked with asterix (*) are due to solvent or change of detector.

clearly exceed typical values for a thermally induced VT transition, which points to partial dissociation.²⁰ The presence of a five-coordinate hs-Co^{II}(SQ)₂ species trans-5 (Scheme 1) was verified by titrating a solution of trans-6 with trans-4-stypy ligand at RT. Upon addition of the ligand, the green colored solution became bluish-green, the 750 nm band decreased and absorption at 600 nm became more developed (Fig. 7). These changes closely resemble those in the variable-temperature experiments (vide supra) and point to a hs-Co^{II}(SQ)₂ \rightarrow ls-Co^{III}(Cat)(SQ) transition. The formation of ls-Co^{III}(Cat)(SQ) species upon titration is further corroborated by the growing IVLLCT band at \sim 2650 nm. An isosbestic point was observed at 1060 nm. Very similar color and spectral changes were observed upon addition of cis-4-stypy ligand to a solution of cis-6 (see $ESI⁺$). Thus, the solution of *cis-*6 contains some five-coordinate cis-5 species. Edge Article.
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Magnetic properties of trans-6 in solution changed on titration as well: the RT magnetic moment decreased gradually from 3.81(6) μ_B (prior to *trans*-4-stypy addition) to 2.84(7) μ_B after addition of trans-4-stypy (10 eq.). This is in agreement with the suggested hs-Co^{II}(SQ)₂ \rightarrow ls-Co^{III}(Cat)(SQ) switching. Similarly to *trans*-6, a decrease in μ_{eff} from 4.1(1) to 3.1(1) μ_{B} was observed upon addition of cis-4-stypy (16 eq.) to a solution of cis-6 (see ESI†). Thus, the VT states in both trans- and cis-complexes can be switched at RT via addition of corresponding pyridine ligands, as we previously documented in Coordination-Induced Valence Tautomerism (CIVT) effect.²⁰

In order to obtain thermodynamic parameters for pure VT transitions between the ls-Co^{III}(Cat)(SQ) and hs-Co^{II}(SQ)₂ redox isomers, we recorded variable-temperature electronic spectra on solutions of trans-6 and cis-6 containing an excess of respective 4-stypy ligand. Thus, the dissociation and formation of five-coordinate species in these solutions must be suppressed. The corresponding van't Hoff fits provided $\Delta H = 39(2)$ kJ mol⁻¹ and $\Delta S = 108(9)$ J mol⁻¹ K⁻¹ for *trans*-6 and $\Delta H =$ 37(2) kJ mol⁻¹ and $\Delta S = 110(7)$ J mol⁻¹ K⁻¹ for cis-6 (see ESI[†]). Note, that these values are smaller than those obtained on solutions without excess of 4-stypy ligands and now they are in

agreement with typical values reported for VT isomers in solution.³⁵

The evolution of the electronic spectrum of trans-6 and the decrease of magnetic moment on titration were both fitted using a non-linear regression approach (see ESI†).³⁹ The intensity of the 750 nm absorption band as a function of trans-4-stypy concentration (Fig. 8) was fitted to yield an association constant $K_a = 4(1) \times 10^3$ L mol⁻¹ (eqn (1)). In excellent agreement with this value, a similar fitting for magnetic moment gave $K_a = 6(2)$ \times 10³ L mol⁻¹ (Fig. 9). Given the values of K_a , the toluene solution of the complex must contain \sim 45% of the five-coordinate *trans*-5 and \sim 55% of the six-coordinate *trans*-6 at given concentrations at RT.

Similarly, the changes of electronic spectrum and magnetic moment of *cis*-6 upon addition of *cis*-4-stypy gave K_a values of $1.1(3) \times 10^3$ and $0.3(1) \times 10^3$ L mol⁻¹, respectively (see ESI†). Although there is some discrepancy in the values obtained by

Fig. 8 The evolution of the 750 nm absorption band of trans-6 dissolved in toluene upon titration with trans-4-stypy at RT. Non-linear regression fit parameters: ε (*trans*-6 = {*trans*-6^{HS} \leftrightarrow *trans*-6^{LS}}) =
1.05/2) \times 10³ M⁻¹ cm⁻¹ e(*trans-*5) = 2.9(1) \times 10³ M⁻¹ cm⁻¹ K = 1.05(2) \times 10³ M⁻¹ cm⁻¹, ε (trans-5) = 2.9(1) \times 10³ M⁻¹ cm⁻¹, $K_a =$
4(1) \times 10³ Lmol⁻¹ $4(1) \times 10^3$ L mol⁻¹.

Fig. 9 The evolution of the effective magnetic moment of trans-6 dissolved in toluene upon titration with trans-4-stypy at RT determined by the Evans method (toluene/[D₈]toluene/TMS = $10:2:1$). Non-linear regression fit parameters: $\mu_{\text{eff}}(trans-6 \equiv \{trans-6^{HS} \leftrightarrow$ trans-6^{LS}}) = 2.51(5) $\mu_{\rm B}$, $\mu_{\rm eff}(trans - 5) = 4.7(2)\mu_{\rm B}$, and $K_{\rm a} = 6(2) \times 10^3$ L mol $^{-1}$.

two methods, it is more important that the K_a values for cis-6 are significantly smaller than those for *trans*-6. Thus, *cis*-4-stypy is a weaker ligand than trans-4-stypy. This is corroborated by X-ray crystallography, showing that trans-4-stypy ligands in trans-6 are planar thus better π -acceptors, than non-planar cis-4-stypy ligands in cis-6.

Note, that the RT magnetic moment of dissolved trans-6 decreased to only $2.8\mu_B$ in the presence of a large excess of trans-4-stypy (20 eq.), but not to the low temperature limit of $1.7\mu_B$. This seeming inconsistency can be readily resolved by considering that the parent trans-6 exists as a mixture of VT isomers trans-6^{LS} \leftrightarrow trans-6^{HS} at RT in solution (eqn (1)). Thus, at low temperatures only *trans*- 6^{LS} is present in solution in agreement with the observed magnetic moment of 1.7 μ B. At RT 45% of trans-6 dissociates to form trans-5 (vide supra). In the presence of a large excess of trans-4-stypy the dissociation is suppressed leaving a mixture of trans- 6^{LS} and trans- 6^{HS} at RT. The estimations from magnetic data yield 78% of $trans-6^{LS}$ and 22% of $trans-6$ ^{HS} present in this solution at RT. The same applies to *cis-*6: in the presence of a large excess of cis-4-stypy the equilibrium in solution at RT is virtually shifted to six-coordinate species cis- 6^{HS} and *cis*- 6^{LS} (eqn (1)). Chemical Science

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5 \xrightarrow{\text{ } 4-4\text{-stypy}} \left\{6^{\text{HS}} \xrightarrow{\text{ } 6^{\text{LS}}} 6^{\text{LS}}\right\} \qquad (1)
$$

Photochemistry

Solutions of *trans*-6 and *cis*-6 differ in transition temperatures $T_{1/2}$ and association constants K_3 . Consequently, the amount of ls-Co^{III}(Cat)(SQ) and hs-Co^{II}(SQ)₂ species in the two solutions is different. Thus, trans \leftrightarrow cis photoisomerization of 4-stypy ligands offers the opportunity to induce a shift in VT equilibrium and to switch magnetic properties at RT.

When *trans*-6 solution was irradiated at $\lambda = 320$ nm, the absorption band at 307 nm decreased in intensity confirming trans

Fig. 10 Changes in absorption spectrum of trans-6 dissolved in toluene upon UV irradiation at RT ($c = 3.7 \times 10^{-4}$ M, $\lambda = 320 \pm 8$ nm, 1000 W Xe lamp). Signals marked with asterix (*) are due to solvent or change of detector.

Fig. 11 Changes in absorption at 750 and 2830 nm of trans-6 dissolved in toluene upon UV irradiation at RT ($c = 3.7 \times 10^{-4}$ M, $\lambda = 320$ \pm 8 nm, 1000 W Xe lamp). The dashed lines serve as guides to the eye.

 \rightarrow *cis* isomerization of 4-stypy (see ESI†). Upon irradiation, the IVLLCT band at \sim 2650 nm decreased and the CT band at 750 nm increased revealing an isosbestic point at 1060 nm (Fig. 10). The photostationary state (PSS) was nearly reached within 60 min (Fig. 11). It is worth mentioning that very similar spectral changes were observed for ls-Co^{III}(Cat)(SQ) \rightarrow hs-Co^{II}(SQ)₂ transition in variable-temperature and CIVT experiments (vide supra). Thus, the *trans* \rightarrow *cis* photoisomerization of 4-stypy did induce a VT transition in solution at RT.

These results are strongly supported by NMR spectroscopy. Upon irradiation, the magnetic moment increased gradually from 3.53(6) to 4.00(6) μ_B and the PSS was reached within 45 min (Fig. 12). Upon further irradiation, no changes in NMR and electronic spectra were detected verifying stability of the PSS. The amount of ls-Co^{III}(Cat)(SQ) species decreased from 60% before irradiation to 43% after irradiation. Thus, the ls-Co^{III}(Cat)(SQ) \rightarrow hs-Co^{II}(SQ)₂ photoconversion of 28% $((60-43)/60 \times 100)$ was achieved at RT. Importantly, the electronic spectrum and magnetic moment remained unchanged upon storing the irradiated solution at RT for at least 20 hours! Thus, the photogenerated VT state reveals extraordinary

Fig. 12 The evolution of magnetic moment of trans-6 dissolved in toluene upon UV irradiation at RT determined by the Evans method $(c = 5.0 \times 10^{-4}$ M, toluene/[D₈]toluene/TMS = 10 : 2 : 1, $\lambda = 320 \pm 8$ nm, 1000 W Xe lamp). The dashed line serves as a guide to the eye.

Fig. 13 The evolution of the X-band EPR spectrum of trans-6 dissolved in benzene upon UV irradiation at RT ($c = 1.0 \times 10^{-4}$ M, $\lambda = 323$ \pm 5 nm, 150 W Xe lamp, frequency: 8.9410 GHz, modulation: 0.7 mT, power: 20 mW). Note, that the signal is due to $Is-Co^{III}(Cat)(SQ)$, whereas hs-Co^{II}(SQ)₂ species are not detected. Relatively noisy signal is due to low concentration of trans-6, required to speed up bulk photolysis, and consequently very small amount of EPR active $\text{ls-Co}^{\text{III}}(\text{Cat})(\text{SQ})$ species present in solution. The spectra of a more concentrated solution and at low temperatures with increased $ls\text{-}Co^{\text{III}}(Cat)(SQ)$ content are given in ESI.†

thermal stability at RT, as compared to common nanoseconds.⁸ This is to expect, since cis-4-stypy shows exceptional high thermal stability.²¹

The ls-Co^{III}(Cat)(SQ) \rightarrow hs-Co^{II}(SQ)₂ photoswitching in trans-6 solution was further confirmed by EPR spectroscopy. The hs-Co^{II}(SQ)₂ isomers are usually EPR silent, whereas the $\text{ls-Co}^{\text{III}}(\text{Cat})(\text{SQ})$ state can be readily detected both at low and RT. Thus, a solution of trans-6 at RT shows a typical isotropic spectrum with $g_{\text{iso}} = 1.9980$ and a ⁵⁹Co super-hyperfine coupling constant $A_{\text{iso}} = 10.7 \times 10^{-4} \text{ cm}^{-1}$, which is synonymous with a ligand-based radical $\text{ls-Co}^{\text{III}}(\text{Cat})(\text{SQ})$.⁴⁰ Upon *in situ* irradiation at $\lambda = 323$ nm at RT, the signal of $\text{ls-Co}^{\text{III}}(\text{Cat})(\text{SQ})$ species gradually decreased, whereas no new signals appeared (Fig. 13). The PSS was reached within 60 min.

We examined the possibility to perform trans \rightarrow cis photoisomerization by low-energy excitation in visible as well.⁴¹ Whereas irradiation at $\lambda = 420$ and 750 nm did not induce desired photoreaction, irradiation at $\lambda = 600$ nm did induce $trans \rightarrow cis$ isomerization. Unfortunately, the latter photoreaction was accompanied by some unidentified side reactions. Thus, the irradiation at $\lambda = 320$ nm is the most efficient and clear way to perform trans \rightarrow cis isomerization in trans-6 solution.

The reverse hs-Co^{II}(SQ)₂ \rightarrow ls-Co^{III}(Cat)(SQ) photoswitching could be accomplished as well starting from cis-6 complex and performing $cis \rightarrow trans$ ligand-based photoisomerization. When cis-6 solution was irradiated at $\lambda = 272$ nm, the cis \rightarrow trans photoisomerization of 4-stypy could be unambiguously confirmed by increasing 306 nm band in electronic spectrum (see ESI†). Simultaneously, the 750 nm CT band decreased in intensity, a shoulder at 600 nm developed, and the LLIVCT band at \sim 2650 nm increased before the PSS was reached within 30 min

Fig. 14 Changes in absorption spectrum of cis-6 dissolved in benzene upon UV irradiation at RT ($c = 5.0 \times 10^{-4}$ M, $\lambda = 272 \pm 8$ nm, 1000 W Xe lamp). Signals marked with asterix (*) are due to solvent or change of detector.

(Fig. 14). The observed spectral changes show exactly opposite trends compared to the *trans* \rightarrow *cis* photoisomerization experiment. Thus, a *cis* \rightarrow *trans* photoisomerization of 4-stypy induces hs-Co^{II}(SQ)₂ \rightarrow ls-Co^{III}(Cat)(SQ) conversion at RT.

The reverse photoswitching of VT states in *cis*-6 solution was further corroborated by the decrease of magnetic moment from 3.9(1) to 3.7(1) μ _B upon irradiation at $\lambda = 272$ nm (Fig. 15). After irradiation the electronic spectrum and magnetic moment remained virtually unchanged during at least 3 hours demonstrating high stability of photogenerated species.

Two mechanisms for the observed photoswitching of VT states should be considered. Upon irradiation at $\lambda = 320$ nm, trans-4-stypy is converted to cis-4-stypy (coordinated and noncoordinated). The non-planar geometry of cis-4-stypy renders it a weaker π -acceptor compared to the *trans*-form. Thus, the ligand field is reduced upon *trans* \rightarrow *cis* isomerization, which should stabilize the hs- $Co^H(SQ)₂$ state in six-coordinate species 6. At the same time, cis-4-stypy is a weaker coordinating ligand than the trans-isomer, 42 as was confirmed by differing associate

Fig. 15 The evolution of magnetic moment of cis-6 dissolved in toluene upon UV irradiation at RT determined by the Evans method $(c = 5.0 \times 10^{-4}$ M, toluene/[D₈]toluene/TMS = 10 : 2 : 1, $\lambda = 272 \pm 8$ nm, 1000 W Xe lamp). The dashed line serves as a guide to the eye.

constants (vide supra). Thus, the equilibrium between fivecoordinate 5 and six-coordinate 6 species is expected to shi upon *trans* \rightarrow *cis* isomerization. In this case some ${6^{LS} \leftrightarrow 6^{HS}}$ species is converted to 5 increasing the hs-Co^{II}(SQ)₂ content (eqn (1)).

Similar considerations apply to reverse reaction, when cis-4 stypy is converted to *trans*-4-stypy by irradiating at $\lambda = 272$ nm. The observed hs-Co^{II}(SQ)₂ \rightarrow ls-Co^{III}(Cat)(SQ) photoconversion may be due to increased ligand field splitting accompanying the \rightarrow *trans* isomerization. Alternatively, the equilibrium between 5 and $\{6^{LS} \leftrightarrow 6^{HS}\}$ may be shifted toward the sixcoordinate species (eqn (1)) due to stronger coordinating trans-4-stypy, thus leading to increased ls- $Co^{III}(Cat)(SQ)$ content.

To gain a better mechanistic understanding, we performed an irradiation experiment on a trans-6 solution containing a large excess of trans-4-stypy (50 eq.), in which the dissociation is virtually suppressed. Upon extended exposure to UV light the absorption spectrum did not show any detectable changes in visible and SWIR regions characteristic for VT conversion. Therefore, we concluded, that the successful photoswitching of VT states in pristine solutions of trans-6 and cis-6 is unlikely due to variation of the ligand field strength, but to great extent due to the light-induced shift in equilibrium between five- and sixcoordinate species. Chemical Science

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It is important to note that the photoswitching of electronic states in trans-6 and cis-6 solutions is of a pure molecular origin. However, the reported irradiation times (typically 30–60 min) correspond the switching of huge number of molecules (bulk photolysis), collectively detected by spectroscopic methods. Since we know how many molecules are switched from ls- $Co^{III}(Cat)(SQ)$ to hs- $Co^{II}(SQ)_{2}$ state upon UV irradiation (appr. 10^{17}), the effective photoswitching rate under our conditions can be estimated as 10^{17} molecules \div 3600 s = 3 \times 10^{13} molecules per s.

In unison with early works by Boillot¹⁵ and recent works of Herges and Tuczek^{42,43} on bistable spin-crossover metal complexes, we coin the observed effect Ligand-Driven Light-Induced Valence Tautomerism (LD-LIVT).

Conclusions

In spite of great efforts on the development of molecular photoswitches on the base of valence tautomeric metal complexes during the last decade, the switching with light has been restricted to very low temperatures (usually below 20 K). Here, for the first time we have demonstrated light-induced switching in valence tautomeric cobalt complexes at room temperature. The driving force for the photoswitching is bidirectional trans \leftrightarrow *cis* photoisomerization of 4-styrylpyridine ligands deliberately integrated into bistable magnetic molecules. The photoswitching results in modulation of magnetic properties in solution at room temperature. The novel effect has been coined Ligand-Driven Light-Induced Valence Tautomerism (LD-LIVT) and proceeds at molecular level with very high effective photoswitching rates. Extensive spectroscopic studies reveal photoconversion of 28% and extraordinary thermal stability of photoinduced states at RT for hours, as compared to common

nanoseconds. Consequently, this work may open new horizons in applications of magnetic switches based on valence tautomeric metal complexes in molecular devices functioning at room temperature.

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Notes and references

- 1 R. M. Buchanan and C. G. Pierpont, J. Am. Chem. Soc., 1980, 102, 4951–4957.
- 2 D. N. Hendrickson and C. G. Pierpont, Top. Curr. Chem., 2004, 234, 63–95.
- 3 J. Sedo, J. Saiz-Poseu, F. Busque and D. Ruiz-Molina, Adv. Mater., 2013, 25, 653–701.
- 4 V. I. Minkin, Russ. Chem. Bull., 2008, 57, 687–717.
- 5 O. Sato, J. Tao and Y.-Z. Zhang, Angew. Chem., Int. Ed., 2007, 46, 2152–2187.
- 6 A. Dei, D. Gatteschi, C. Sangregorio and L. Sorace, Acc. Chem. Res., 2004, 37, 827–835.
- 7 T. Tezgerevska, K. G. Alley and C. Boskovic, Coord. Chem. Rev., 2014, 268, 23–40.
- 8 D. M. Adams, B. L. Li, J. D. Simon and D. N. Hendrickson, Angew. Chem., Int. Ed., 1995, 34, 1481–1483.
- 9 O. Sato, S. Hayami, Z.-Z. Gu, K. Seki, R. Nakajima and A. Fujishima, Chem. Lett., 2001, 30, 874–875.
- 10 F. Varret, M. Nogues and A. Goujon, in Magnetism: Molecules to Materials, ed. J. S. Miller and M. Drillon, Wiley-VCH Verlag, New York, 2001, pp. 257–295.
- 11 O. Sato, A. L. Cui, R. Matsuda, J. Tao and S. Hayami, Acc. Chem. Res., 2007, 40, 361–369.
- 12 A. Beni, C. Carbonera, A. Dei, J. F. Letard, R. Righini, C. Sangregorio and L. Sorace, J. Braz. Chem. Soc., 2006, 17, 1522–1533.
- 13 R. D. Schmidt, D. A. Shultz, J. D. Martin and P. D. Boyle, J. Am. Chem. Soc., 2010, 132, 6261–6273.
- 14 R. D. Schmidt, D. A. Shultz and J. D. Martin, Inorg. Chem., 2010, 49, 3162–3168.
- 15 M.-L. Boillot, J. Zarembowitch and A. Sour, Top. Curr. Chem., 2004, 234, 261–276.
- 16 Molecular Switches, ed. B. L. Feringa and W. R. Browne, WILEY-VCH Verlag & Co. KGaA, Weinheim, Germany, 2011.
- 17 M. M. Paquette, R. A. Kopelman, E. Beitler and N. L. Frank, Chem. Commun., 2009, 5424–5426.
- 18 M. Milek, A. Witt, C. Streb, F. W. Heinemann and M. M. Khusniyarov, Dalton Trans., 2013, 42, 5237–5241.
- 19 M. Milek, F. W. Heinemann and M. M. Khusniyarov, Inorg. Chem., 2013, 52, 11585–11592.
- 20 A. Witt, F. W. Heinemann, S. Sproules and M. M. Khusniyarov, Chem.–Eur. J., 2014, 20, 11149–11162.
- 21 J. L. R. Williams, R. E. Adel, J. M. Carlson, G. A. Reynolds, D. G. Borden and J. A. Ford, J. Org. Chem., 1963, 28, 387–390.
- 22 R. M. Buchanan, B. Fitzgerald and C. G. Pierpont, Inorg. Chem., 1979, 18, 3439–3444.
- 23 M.-C. Chiang and W. Hartung, J. Org. Chem., 1945, 10, 21–25.
- 24 E. Bill and JulX, version 1.5, MPI for Bioinorganic Chemistry, Muelheim/Ruhr, Germany, 2008.
- 25 F. Neese, Diploma thesis, University of Konstanz, Konstanz, Germany, 1993.
- 26 D. F. Evans, J. Chem. Soc., 1959, 2003–2005.
- 27 SADABS 2008/1, Bruker AXS, Inc., Madison, WI, 2009.
- 28 G. Sheldrick, Acta Crystallogr., Sect. A: Found. Crystallogr., 2008, 64, 112–122.
- 29 CCDC 1015850–1015852 and 1058866 contain the supplementary crystallographic data for this paper.
- 30 C. G. Pierpont, Coord. Chem. Rev., 2001, 216, 99–125.
- 31 C. Roux, J. Zarembowitch, B. Gallois, T. Granier and R. Claude, Inorg. Chem., 1994, 33, 2273–2279.
- 32 M. L. Boillot, S. Pillet, A. Tissot, E. Rivière, N. Claiser and C. Lecomte, Inorg. Chem., 2009, 48, 4729–4736.
- 33 Variation of the high-temperature limit of the effective magnetic moment between 5.0 and 6.0 μ _B does not change the fit parameters significantly (see ESI†).
- 34 Y. Mulyana, G. Poneti, B. Moubaraki, K. S. Murray, B. F. Abrahams, L. Sorace and C. Boskovic, Dalton Trans., 2010, 39, 4757–4767.
- 35 D. M. Adams and D. N. Hendrickson, J. Am. Chem. Soc., 1996, 118, 11515–11528.
- 36 H. Rau, in Photochromism. Molecules and Systems, ed. H. Dürr and H. Bouas-Laurent, Elsevier, Amsterdam, 2003, pp. 64– 164.
- 37 N. A. Pavlova, A. I. Poddel'sky, A. S. Bogomyakov, G. K. Fukin, V. K. Cherkasov and G. A. Abakumov, Inorg. Chem. Commun., 2011, 14, 1661–1664.
- 38 N. S. Hush, in Progress in Inorganic Chemistry, John Wiley & Sons, Inc., 1967, pp. 391–444.
- 39 Note, that the non-linear regression fit used to obtain association constants implies only two species involved into equilibrium, whereas actually three species $(5, 6^{LT}$ and 6^{HT} are present in solution. However, the employed model is still valid, if the temperature is kept constant and thus the ratio between 6^{LT} and 6^{HT} remains constant too. Hence, the model describing the equilibrium between 5 and a quasi-species $\{6^{\text{LT}} \leftrightarrow 6^{\text{HT}}\}$ is correct. Edge Article

20 A. Witt, T. W. Heinemann, S. Sprooks and 35 D.M. Administration. J. Article. Science. 2004, 10:40:44 AM. This article is licensed under a Creative Commons Access Article is licensed under a Creative Commo
	- 40 D. M. Adams, L. Noodleman and D. N. Hendrickson, Inorg. Chem., 1997, 36, 3966–3984.
	- 41 A. Tissot, M.-L. Boillot, S. Pillet, E. Codjovi, K. Boukheddaden and L. M. L. Daku, J. Phys. Chem. C, 2010, 114, 21715–21722.
	- 42 S. Thies, H. Sell, C. Schütt, C. Bornholdt, C. Näther, F. Tuczek and R. Herges, J. Am. Chem. Soc., 2011, 133, 16243–16250.
	- 43 S. Venkataramani, U. Jana, M. Dommaschk, F. D. Sonnichsen, F. Tuczek and R. Herges, Science, 2011, 331, 445–448.